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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO
10/772,963	02/05/2004	David P. Bingaman	2471 US	5299
7590 06/15/2005			EXAMINER	
Teresa J. Schultz			HUI, SAN MING R	
Alcon Research, Ltd. 6201 South Freeway, Q-148			ART UNIT	PAPER NUMBER
Fort Worth, TX 76124-2099			1617	

DATE MAILED: 06/15/2005

Please find below and/or attached an Office communication concerning this application or proceeding.

		Application No.	Applicant(s)			
Office Action Summary		10/772,963	BINGAMAN ET AL.			
		Examiner	Art Unit			
		San-ming Hui	1617			
The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply						
THE MAILING C - Extensions of time r after SIX (6) MONTI - If the period for reply - If NO period for repl - Failure to reply with Any reply received b	DATE OF THIS COMMUNICATION THE AVAILABLE OF THIS COMMUNICATION OF THE AVAILABLE OF THE AVAI	R 1.136(a). In no event, however, may a reply be t	imely filed ays will be considered timely. m the mailing date of this communication. ED (35 U.S.C. § 133).			
Status						
1)⊠ Responsi	ve to communication(s) filed on 0	9 March 2005.				
2a) ☐ This action	• • • • • • • • • • • • • • • • • • • •	This action is non-final.				
3)☐ Since this	·					
Disposition of Clai	ms					
4a) Of the 5) ☐ Claim(s) _ 6) ☑ Claim(s) _ 7) ☐ Claim(s) _	# 1-18 is/are pending in the applicate above claim(s) is/are with is/are allowed. # 1-18 is/are rejected. # 1-18 is/are objected to. # are subject to restriction are	drawn from consideration.				
Application Papers	3		·			
9)☐ The specification is objected to by the Examiner. 10)☐ The drawing(s) filed on is/are: a)☐ accepted or b)☐ objected to by the Examiner.						
Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a).						
Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).						
11) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152.						
Priority under 35 U	l.S.C. § 119					
 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: 1. Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No. 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 						
Attachment(s) 1) Notice of Reference	ces Cited (PTO-892)	4) 🔲 Interview Summar	ry (PTO-413)			
2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08) Paper No(s)/Mail Date 2-3-05. Pager No(s)/Mail Date 2-3-05.						

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DETAILED ACTION

Applicant's amendments filed March 9, 2005 have been entered. The addition of claims 3-18 is acknowledged. Claims 1-18 are pending.

The outstanding rejection under 35 USC 102(b) is withdrawn in view of the applicant's remarks that anecortave acetate as not a glucocorticoid.

The outstanding double patenting rejection is also withdrawn in view of the fact that anecortave acetate is not a glucocorticoid.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

Claims 1-5 and 8-18 are rejected under 35 U.S.C. 103(a) as being unpatentable over Peyman et al. (US patent 5,516,522) and Clark, Clark is reference of record.

Peyman teaches prednisolone, prednisolone acetate, triamcinolne, fluoromethalone, and fluoromethalone acetate as useful in treating proliferative vitreoretinopathy (PVR), an ocular angiogenesis-associated disorder (See col. 7, lines 33-55, especially lines 50, 51, 54). Peyman also teaches the ocular formulation may be as intraocular implant (See the abstract and claim 1).

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Clark teaches anecortave acetate as useful in treating ocular neovascularization condition (See claims 1-5). Clark also teaches the composition can be formulated and administered as intraocular injection (See col. 4, lines 50).

The references taken together do not expressly teach the incorporation of both the herein claimed steroids and anecortave acetate together in a method of treating angiogenesis disorder such as PVR. The references taken together do not expressly teach the herein claimed dosages.

It would have been obvious to one of ordinary skill in the art at the time the invention was made to incorporate the herein claimed steroids and anecortave acetate together in a method of treating angiogenesis disorder such as PVR. It would have been obvious to one of ordinary skill in the art at the time the invention was made to employ the herein claimed dosage to treat PVR.

One of ordinary skill in the art would have been motivated to incorporate the herein claimed steroids and anecortave acetate together in a method of treating angiogenesis disorder such as PVR since the agents are well-known to be useful in treating PVR or neovascularization individually. Therefore, concomitantly employing both agents in a method for the same indications would be *prima facie* obvious (See *In re Kerkhoven* 205 USPQ 1069).

Furthermore, one of ordinary skill in the art would have been motivated to employ the herein claimed dosage to treat PVR since the optimization of result effect parameters (dosage range, dosing regimens) is obvious as being within the skill of the artisan.

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Claims 1-2, 4-5, and 16-18 are rejected under 35 U.S.C. 103(a) as being unpatentable over WO95/03807('807) and Clark.

'807 teaches a method of treating neovascular macular degeneration, an ocular angiogenesis disorder, by administration of triamcinolne (See the abstract, claims 22-25). '807 teaches the routes of administration may be intravitreal injection (See page 3, lines 19-25).

Clark teaches anecortave acetate as useful in treating ocular neovascularization condition (See claims 1-5). Clark also teaches the composition can be formulated and administered as intraocular injection (See col. 4, lines 50).

The references taken together do not expressly teach the incorporation of both the triamcinolone and anecortave acetate together in a method of treating angiogenesis disorder such as neovascular macular degeneration. The references taken together do not expressly teach the herein claimed dosages.

It would have been obvious to one of ordinary skill in the art at the time the invention was made to incorporate triamcinolone and anecortave acetate together in a method of treating angiogenesis disorder such as neovascular macular degeneration. It would have been obvious to one of ordinary skill in the art at the time the invention was made to employ the herein claimed dosage to treat neovascular macular degeneration.

One of ordinary skill in the art would have been motivated to incorporate triamcinolone and anecortave acetate together in a method of treating angiogenesis disorder such as neovascular macular degeneration since the agents are well-known to

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be useful in treating neovascular macular degeneration individually. Therefore, concomitantly employing both agents in a method for the same indications would be *prima facie* obvious (See *In re Kerkhoven* 205 USPQ 1069).

Furthermore, one of ordinary skill in the art would have been motivated to employ the herein claimed dosage to treat neovascular macular degeneration since the optimization of result effect parameters (dosage range, dosing regimens) is obvious as being within the skill of the artisan.

Claims 1-3 and 6-7 are rejected under 35 U.S.C. 103(a) as being unpatentable over Clark and US 4,686,214 ('214).

Clark teaches anecortave acetate as useful in treating ocular neovascularization inflammatory condition (See claims 1-5). Clark also teaches the composition can be formulated and administered as intraocular injection (See col. 4, lines 50).

'214 teaches rimexolone as useful in treating ocular inflammation (See claim 2).

The effective dosage of rimexolone taught as 0.05 to 2.0% (See col. 2, line 59-60).

The references taken together do not expressly teach the incorporation of both rimexolone and anecortave acetate together in a method of treating angiogenesis inflammatory disorder. The references taken together do not expressly teach the herein claimed dosages.

It would have been obvious to one of ordinary skill in the art at the time the invention was made to the incorporation of both rimexolone and anecortave acetate together in a method of treating angiogenesis inflammatory disorder. It would have

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been obvious to one of ordinary skill in the art at the time the invention was made to employ the herein claimed dosage to treat angiogenesis inflammatory disorder.

One of ordinary skill in the art would have been motivated to the incorporation of both rimexolone and anecortave acetate together in a method of treating angiogenesis inflammatory disorder since the agents are well-known to be useful in treating ocular inflammation individually. Therefore, concomitantly employing both agents in a method for treating ocular inflammation associated with angiogenesis would be *prima facie* obvious (See *In re Kerkhoven* 205 USPQ 1069).

Furthermore, one of ordinary skill in the art would have been motivated to employ the herein claimed dosage to treat ocular inflammation associated with angiogenesis since the optimization of result effect parameters (dosage range, dosing regimens) is obvious as being within the skill of the artisan.

Response to Arguments

Applicant's arguments with respect to claims 1-18 have been considered but are moot in view of the new ground(s) of rejection.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to San-ming Hui whose telephone number is (571) 272-0626. The examiner can normally be reached on Mon 9:00 to 1:00, Tu - Fri from 9:00 to 6:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Sreeni Padmanabhan, PhD., can be reached on (571) 272-0629. The fax

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phone number for the organization where this application or proceeding is assigned is 703-872-9306.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

San-ming Hui Primary Examiner

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